AMENDMENTS TO THE CLAIMS

This listing of claims replaces all previous versions, and listings, of claims pending in this

application.

Listing of Claims

1-9. (Canceled)

10. (Currently amended) A method for synthesizing a peptide dimer, comprising:

(a) providing first and second peptide chains linked to a linking moiety L_K, said chains

each possessing multiple amino acid residues capable of disulfide bond formation upon

oxidation; and

(b) oxidizing in a single oxidation step said peptide chains in a manner effective to

preferentially promote formation of disulfide bonds between residues in the same peptide chain

relative to formation of disulfide bonds in different peptide chains, and wherein at least 50% of

said peptide dimer comprises a peptide chain having an intrapeptide disulfide bond.

11. (Previously presented) The method of claim 10, wherein step (b) comprises treatment

with an oxidizing composition containing an oxidizing reagent in an amount effective to

minimize reaction products in which a residue of the first peptide chain binds to a residue of the

second peptide chain.

12. (Previously presented) The method of claim 11, wherein the oxidizing reagent is

dimethyl sulfoxide.

13. (Previously presented) The method of claim 12, wherein the oxidizing composition

comprises approximately 15% to 100% (v/v) dimethyl sulfoxide.

14. (Previously presented) The method of claim 13, wherein the oxidizing composition

comprises approximately 50% to 100% (v/v) dimethyl sulfoxide.

15. (Canceled)

2

Preliminary Amendment in Response to Final Office Action (Dated: November 8, 2007 – Paper No. 20071107)

Application Serial No. 10/737,245

Application Serial No. 10/757,243

Attorney's Docket No. 44368-0005 C1

16. (Previously presented) The method of claim 14, wherein the oxidizing composition comprises approximately 80% to 100% (v/v) dimethyl sulfoxide.

17. (Previously presented) The method of claim 16, wherein the oxidizing composition

comprises approximately 100% (v/v) dimethyl sulfoxide.

18. (Currently amended) The method of claim 10, claim 1, wherein the first peptide chain

is approximately 10 to 40 amino acid residues in length, binds to the erythropoietin receptor, and

contains a sequence of amino acids X3X4X5GPX6TX7X8X9, wherein X3 is C or homocysteine

(Hoc), X4 is R, H, L or W, X5 is M, F, I or nor-leucine (J), X6 is selected from any one of the 20

conventional amino acids and J, X7 is W, 1-naphthylalanine (B) or 2-naphthylalanine (U), X8 is

D, E, I, L, or V, and X9 is C or Hoc; and the second peptide chain is approximately 10 to 40

amino acid residues in length, binds to the erythropoietin receptor, and contains a sequence of

amino acids X'3X'4X'5X'6X'7X'8X'9, wherein X'3 is C or Hoc, X'4 is R, H, L or W, X'5 is M, F,

I or J, 1, X'6 is selected from any one of the 20 conventional amino acids and J, X'7 is W, B or U,

X'8 is D, E, I, L or V,- and X'9 is C or Hoc.

19. (Previously presented) The method of claim 18, wherein one or more of said amino

acid residues are genetically coded L-amino acids.

20. (Previously presented) The method of claim 18, wherein the amino terminus of at

least one of said peptide chains is modified.

21. (Previously presented) The method of claim 10, wherein at least one of said peptide

chains comprises a non-naturally occurring amino acid residue.

22. (Previously presented) The method of claim 10, wherein at least one of said peptide

chains comprises an amino acid residue, wherein a naturally occurring side chain of said amino

acid residue is replaced with a non-naturally occurring side chain.

23. (Previously presented) The method of claim 10, wherein said first peptide chain binds

to the erythropoietin receptor and wherein said second peptide chain binds to the erythropoietin

receptor.

3

Preliminary Amendment in Response to Final Office Action (Dated: November 8, 2007 – Paper No. 20071107)

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Attorney's Docket No. 44368-0005 C1